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- (71) Applicants: THE UNIVERSITY OF PENNSYLVANIA [US/US]; Center for Technology Transfer, Suite 300, 3700 Market Street, Philadelphia, PA 19104-3147 (US). WAYNE STATE UNIVERSITY [US/US]; Technology Transfer Office, 4032 Faculty/Administration Building, 656 West Kirby, Detroit, MI 48202 (US).
- (72) Inventors: SCHREIBER, Alan, D.; The University of Pennsilvania, Centre for Technology Transfer, Suite 300, 3700 Market Street, Philadelphia, PA 19104-3147 (US). WORTH, Randall; The University of Pennsylvania, Centre for Technology Transfer, Suite 300, 3700 Market Street, Philadelphia, PA 19104-3147 (US). PETTY, Howard, R.; Wayne Stete University, Transfer Technology Office, 4032 Faculty/Administration Building, 656 West Kirby, Detroit, MI 48202 (US).

- (74) Agent: WILSON, Mary, J.; Nixon & Vanderhye P.C., 1100 North Glebe Road, Suite 800, Arlington, VA 22201-4714 (US).
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(54) Title: TRIPEPTIDE OF FCγRIIA

(57) Abstract: The present invention relates, in general, to phagocytosis and phagolysosomal fusion and, in particular, to a tripeptide of FeyRIIA that mediates trafficking of targets phagocytosed via FeyRIIA to the lysosomal compartment.

TRIPEPTIDE OF FCYRIIA

This application claims priority from Provisional Application No. 60/252,460, filed November 22, 2000, the entire content of which is 5 incorporated herein by reference.

TECHNICAL FIELD

The present invention relates, in general, to phagocytosis and phagolysosomal fusion and, in particular, to a tripeptide of FcyRIIA that mediates trafficking of targets phagocytosed via FcYRIIA to the lysosomal compartment.

BACKGROUND

Phagolysosome fusion is an important pathway in the degradation of internalized particles. Once a particle is internalized by phagocytosis it is directed toward the lysosomal compartment for degradation. Various studies have traced this sequence of events from binding and phagocytosis to eventual trafficking to lysosomes. In addition, the signaling machinery needed to perform many of these activities has been described. Recently, intracellular tyrosine-based activation motifs (ITAM) have taken center stage in the initiation and propagation of activation signals of phagocytic receptors. 25

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ITAM motifs contribute to the ability of phagocytic receptors to efficiently internalize particles (Tuijnman et al, Blood 79:1651 (1992), Mitchell et al, Blood 84:1753 (1994)). ITAM motifs 5 are composed of two YXXL motifs separated by a string of various amino acids. This motif forms a SH-2 binding domain for docking of signaling proteins such as Src and Syk, among others (Isakov Immunol. Res. 16:85 (1997), Isakov, J. Leuko. Biol. 10 61:6 (1997)). Specifically, upon ITAM phosphorylation, FcYRIIA has been shown to signal through Syk (Indik, et al, Blood 86:4389 (1995), Matsuda et al, Mol. Bio. Cell 7:1095 (1996)). In addition, mutation of either of the ITAM tyrosines abolishes the phagocytic activity of FcyRIIA 15 (Mitchell et al, Blood 84:1753 (1994)). These YXXL

Once a target is internalized, it can be sent to the lysosomal compartment for degradation. Dileucine motifs in the cytoplasmic domain of various receptors are responsible for the trafficking of targets from phagosomes to lysosomes (Mayorga et al, J. Biol. Chem. 266:6511 (1991), Hunziker and Fumey, EMBO J. 13:2963 (1994), Letournier and Klausner,

sequences can also associate with adaptor proteins

such as AP-1 and AP-2 in forming clathrin cups

during phagocytosis.

Cell 69:1143 (1992)). This motif is present in many receptors such as FcyRIIB, the LDL receptor, and the mannose 6-phosphate receptor (Matter et al, J. Cell

30 Biol. 126:991 (1994), Johnson et al, J. Biol. Chem.

267:17110 (1992)). Mutation of either or both of the leucine residues in these receptors significantly reduces or abolishes lysosomal delivery, respectively.

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FCYRIIA mediates phagocytosis through an ITAM motif and also mediates phagolysosomal fusion (Mitchell et al, Blood 84:1753 (1994)). However, there is no consensus di-leucine motif located in the cytoplasmic domain of FCYRIIA. Therefore, another sequence in the cytoplasmic domain of FCYRIIA must participate in lysosomal trafficking. The present invention relates to that sequence.

SUMMARY OF THE INVENTION

The present invention relates to a tripeptide

of FcqRIIA that mediates trafficking of targets

phagocytosed via FcqRIIA to the lysosomal

compartment.

Objects and advantages of the present invention will be clear from the description that follows.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1. A distinct FcyRIIA cytoplasmic domain sequence determines phagolysosomal fusion. CHO cells were transfected with WT FcyRIIA (WT IIA, column 2) or with mutants of the FcyRIIA cytoplasmic ITAM. Wt IIA contains the ITAM sequence Y2MTL-Y3LTL. The FcyRIIA mutants contain the following ITAM sequences: Y2MTL-Y3ATL (designated Y3ATL,

column 3), Y2MTL-Y3LTA (designated Y3LTA, column 4), Y2MTL-Y3ATA (designated Y3ATA, column 5) or F2MTL-F3LTL (designated Y2FY3F, column 6) (Y=tyrosine, M=methionine, T=threonine, L=leucine, A=alanine, F=phenylalanine). After 48 hrs, the transfected cells were loaded with rhodamine conjugated dextran and then incubated with IgG coated RBCs (EA). Following removal of externally bound EA, the phagocytic index (PI), the number of internalized EA/100 cells, was determined by bright field microscopy. Lysosomes labelled with rhodamine conjugated dextran were visualized by fluorescence microscopy. Phagolysosome fusion was analyzed by determining the co-localization of EA and rhodamine dextran and expressed as % co-localization. 15 Column 1 represents sham transfected cells.

Mutation of either or both leucines in the Y3LTL sequence of the FcyRIIA ITAM inhibits phagolysosomal fusion but does not inhibit phagocytosis of EA. It has been previously demonstrated that FcyRIIA in the absence of ITAM tyrosines (Y2FY3F) does not mediate phagocytosis. However, phagocytosis of EA is partially restored for Y2FY3F by co-transfection with the complement receptor type 3 (CR3) (Worth et al, J. Immunol. 157:5660-5665 (1996)) as demonstrated in column 6. In co-transfected cells, Y2FY3F and CR3 interact and EA bound to Y2FY3F are phagocytosed through the cytoplasmic domain of CR3. 78% of the ingested EA mediated by CR3 and Y2FY3F co-localized with

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lysosomes (column 6), indicating that the ITAM tyrosines do not play a significant role in phagolysosomal fusion. Significant inhibition of phagolysosomal fusion (p<.001) was observed for the mutants Y3ATL, Y3LTA and Y3ATA, while the ingestion of EA (phagocytosis) was unaltered (columns 3-5). Thus the LTL sequence of the FcγRIIA cytoplasmic domain targets the phagosome for fusion with lysosomes whereas the tyrosines of the ITAM sequence are essential for the initial stage of phagocytosis.

Figure 2. Mutation of the novel L-T-L motif in the cytoplasmic domain of FcyRIIA inhibits phagolysosome fusion.

Figure 3. L-T-L motif mediates specific targeting of internalized targets to fuse with lysosomes.

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Figure 4. L-T-L motif inhibits fusion events leading to phagolysosome formation but not protein colocalization.

Figure 5. Inserting the L-T-L motif into a receptor that normally does not mediate efficient phagolysosome formation increases the ability to form phagolysosomes.

DETAILED DESCRIPTION OF THE INVENTION

The present invention is based on the realization that the cytoplasmic domain of FcYRIIA mediates lysosome fusion subsequent to phagocytosis. This L-T-L motif is found at the C-terminal of the ITAM motif of FcYRIIA.

Chinese hamster ovary (CHO) cells provide a good model system for studying phagocytosis and intracellular trafficking. CHO cells transiently transfected with FcyRIIA bind and internalize IgG-coated targets efficiently. Internal targets can be differentiated from bound targets by the addition of a fluorescent secondary goat anti-rabbit IgG. The second step antibody binds only to bound targets thus discriminating between bound and internal targets. In addition, FcyRIIA has been shown to mediate lysosomal fusion by observing the co-localization of pre-loaded fluorescent dextran, which accumulates in lysosomes, with the target when viewed with fluorescence microscopy.

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FcγRIIA mediated lysosome fusion does not require an intact ITAM motif. Previously described studies have shown that mutation of either of the tyrosine residues in the ITAM motif of FcγRIIA abolishes phagocytosis (Mitchell et al, Blood 84:1753 (1994)). Because mutation of the two tyrosine residues abolishes phagocytosis, the genetic complementation ability of complement receptor type 3 (CR3) was utilized. CR3 has

previously been shown to rescue the phagocytic activity of mutated FcqRIIA (Worth et al, J. Immunol. 157:5660 (1996)). FcqRIIA with mutations of tyr—phe is able to mediate lysosomal delivery of targets phagocytosed through the complementary activity of CR3 (which itself does not mediate lysosomal fusion). Mutation of the two tyrosine residues comprising the ITAM abolishes the phagocytic activity of FcqRIIA. However, in the presence of CR3, phagocytosis is restored and over 90% of those internalized targets are delivered to lysosomes. These data indicate that lysosomal delivery is a distinctly separate signal from that involved in phagocytosis and is not dependent on an active ITAM motif.

That lysosomal trafficking and phagocytosis are separate signals is confirmed by mutating the L-T-L sequence of FcyRIIA and observing the ability of internalized targets to be delivered to lysosomes. Firstly, mutation of any or all of these residues 20 does not significantly affect the phagocytic activity of FcyRIIA. Secondly, mutation of either or both of the leucine residues effectively inhibits 70% of internalized particles from fusing with lysosomes. In addition, mutation of the threonine 25 residue alone reduces the lysosomal targeting capacity of FcyRIIA by nearly 70%. However, mutation of all three of these residues does not affect phagocytosis but decreases the lysosomal

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delivery ability to that of a tailless mutant FCYRIIA.

A similar receptor was also studied that does not contain a di-leucine or L-T-L motif. The γ 5 .chain utilized by various Fc receptors such as FcyRI and FcyRIIIA, was utilized. A chimeric FcyRIIIA was formed containing the extracellular domain of FCYRIIIA and the transmembrane and cytoplasmic domains of the y chain. This chimeric receptor containing the \gamma chain signaling machinery is not 10 able to target internalized phagocytosed particles to lysosomes. The ITAM motif of the γ chain was then mutated to contain a L-T-L motif and lysosomal delivery ability studied (Fig. 5). In the presence of the L-T-L motif, the γ chain is able to target internalized particles to lysosomes. This study shows that lysosome targeting ability can be transferred to other receptors by translocating this L-T-L motif.

The L-T-L motif in the cytoplasmic domain of FCYRIIA thus mediates lysosome fusion. FCYRIIAmediated phagocytosis and lysosomal trafficking are composed of two distinct steps mediated by individual signaling motifs. Separate and distinct signals used to mediate internalization and targeting has previously been proposed for the CD3 chains of the T-cell receptor (Letourneur and Klausner, Cell 69:1143 (1992)). The studies described herein confirm that these signals can be 30 distinct, independently acting moieties.

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activities of various secondary signal molecules such as Syk, Rac, Rab, and Rho have all been implicated in endosomal/lysosomal dynamics. Further studies are needed to show which signaling molecules are required for various steps of the internalization pathway. These activities may involve a relay type interaction whereby upon receptor activation by phosphorylation, Syk or another kinase can bind. Once Syk is released, the signal may propagate further by activation of Rac/Rab/Rho or another molecule directing the particle to the lysosomal compartment.

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The demonstration that the L-T-L motif in the cytoplasmic domain of FcyRIIA is responsible for mediating phagolysosomal fusion makes possible gene therapy strategies whereby a sequence encoding naturally occurring FcyRIIA or a modified form of FcyRIIA (e.g., a form modified so as to include more than one L-T-L motif (e.g., 2 or 3 L-T-L motifs) is transferred into target cells that either normally express FcyRIIA or cells that do not normally express FcyRIIA but that can be effective in cleaning, for example, bacterial infections. Examples of target cells include endothelial cells, fibroblasts, macrophage and epithelial cells (such as hepatocytes and bronchial epithelial cells). The receptor encoding sequence can be administered as naked DNA, in a liposome or bacterium or it can be present in a vector, e.g., a viral vector such an

adenoviral or adenoassociated vector or a retroviral vector.

The demonstration that the L-T-L motif in the cytoplasmic domain of FcqRIIA is responsible for mediating phagolysosomal fusion also makes it possible to alter the sequences of Fc receptors, naturally incapable of mediating phagolysosomal fusion, so that they possess that activity.

Transferring the L-T-L motif to such receptors

(e.g., receptors for mycobacterium including CR3 toll-like receptors, etc.) can increase the efficiency of bacterial killing. Sequences encoding such receptors can be used in gene therapy regimens, as described above.

More specifically, the Fc receptor γ chain by itself does not efficiently mediate phagolysosomal fusion; however, when an L-T-L sequence is inserted into its cytoplasmic domain, it mediates phagolysosomal fusion with increased efficiency
(Figure 5). Thus, in such a manner, receptors that do not mediate phagolysosomal fusion can be induced to do so. As indicated above, Fc receptors can also be altered so as to enhance their natural ability to mediate phagolysosomal fusion. For example,

FcγRIIA, upon addition of further L-T-L sequences to the cytoplasmic domain, becomes more potent and efficient in mediating phagolysosomal fusion.

(Increasing the number of FcγRIIA molecules (e.g., by administering a biologically active molecule) can be used as an alternative means of increasing the

number of L-T-L sequences.) These approaches can be usefully applied for enhancing the killing of bacteria, fungi and other microorganisms (e.g., pyrogenic bacteria such as E. coli, S. aureus and P. aeruginosa). Some microorganism survive intracellularly, such as mycobacterium, leishmania and listeria. Enhancing phagolysosomal fusion of these antibody coated microorganisms is useful in controlling the growth and killing of these 10 microorganisms.

In addition to mycobacterium, fungi and other bacteria, the anthrax bacterium can also be targeted to increase the efficiency of its (B. anthracis) being killed. For example, the uptake of anthrax spores by FcyRIIA or another cell receptor can be induced to undergo phagolysosomal fusion.

The demonstration that the L-T-L motif is responsible for mediating phagolysosomal fusion also makes apparent the advantage of targeting microbes to FcyRIIA using therapeutic strategies involving, for example, the use of a bi-specific antibody that recognizes the target microbe and the extracellular domain of FcyRIIA specifically.

The invention further relates to methods of inducing FcyRIIA uptake and targeting of a microorganism or other particle (e.g., an immune complex) to phagolysosomes of macrophages and other leukocytes. In accordance with this method, IgG antibody directed at the microorganism (e.g.,

30 bacterium, including antibiotic resistant E. coli,

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Staphylococcus, etc, mycobacterium, anthrax bacterium, (e.g., Bacillus anthracis or B. anthracis spores) is administered. The antibody used can be an IgG antibody that recognizes the microorganism and that has associated therewith an L-T-L containing peptide.

The invention also relates to methods of increasing the number of Fc γ RIIA molecules per macrophage or leukocyte. Certain small molecules can be used to effect the increase, as can IFN- γ or IL-4 inhibitor or cytokine that inhibits the release or action of IL-4.

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The invention further relates to a method of facilitating targeting of a microorganism to the phagolysosome by administering an L-T-L containing peptide in a manner such that it associates with the microorganism. A liposome containing or otherwise associated with an L-T-L containing peptide can be injected IV or in some other manner such that it is targeted to macrophages/leucocytes.

In accordance with the invention, Fc receptors naturally capable of mediating phagolysosomal fusion can be rendered incapable of such mediation. This can be accomplished using peptide mimetics or small molecule (organic) mimetics that function as inhibitors of the L-T-L sequence. This approach is advantageous when it is desirable to inhibit phagolysosomal fusion, for example, in the preservation of antibody and immune complexes and in hindering their degradation.

Receptors as indicated above (including modified receptors) and L-T-L containing peptides (and respective encoding sequences) can be administered using techniques described, for example, in USP 6,608,983, 5,858,981, 5,821,071, 5,776,910, 5,641,875, and 5,641,863.

Certain aspects of the invention can be described in greater detail in the non-limiting Examples that follows. The following references

10 include details of the receptor structures and encoding sequences: Schreiber et al, Clin. Immunol. Immunopath. 62:S66 (1992), Cassel et al, Molec. Immunol. 30:451 (1993), Allen et al, Science 243:378 (1989), Letourner et al, J. Immunol. 147:2652

15 (1991), Ra et al, Nature (Lond.) 241:752 (1989), Park et al, Clin. Res. 41:324A (1993), Simmons et al, Nature 333:568 (1988).

EXAMPLE 1

Cytoplasmic Domain of FcyRIIA (CD32).
Participates in Phagolysosomal Formation

Experimental Details

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· Cell culture and transfections

Chinese hamster ovary (CHO) cells were transfected by electroporation with a mixture of 1.5 μ g of pSVneo, 5 μ g of pBACD11b (generated by replacing the CD11a cDNA in pBACD11a (Krauss, Hum. Gene Ther. 2:221 (1991)) with the CD11b cDNA (Hickstein et al, Proc. Natl. Acad. Sci. USA 86:257 (1989)), 5 μ g of pCMVBACD18, and 5 μ g of either

pRcCMVCD32 or a variant of this CD32 plasmid containing a tail-minus mutation, as described (Xue, et al, J. Immunol. 152:4630 (1994)). Expansion and selection were performed as previously described 5 (Worth et al, J. Immunol. 157:5660 (1996)). Six different clones were generated: 161-24 which was not transfected but exposed to the transfection protocol; 161-84 expressed only CR3; 131-3 which expressed wild type FcyRIIA; 135-12 expresses FCYRIIA tailless alone; 169-8, 169-24 which both express the FcyRIIA tailless and the α and β chains of CR3; and 173-46 expressing both the wild-type FcyRIIA and CR3. In addition, variants expressing a full length FcyRIIA cytoplasmic domain with tyrosine-phenylalanine mutations in both of the ITAM motifs (FcYRIIA ITAM mutant) were transiently transfected into an untransfected CHO cell line (labeled clone 161-30) or a CR3 expressing CHO cell line (labeled clone 169-85) using DEAE dextran or 20 FuGene6 transfection reagent. For experiments, cells were seeded onto 25mm² coverslips and allowed to adhere overnight at 370C in 5% CO2. Cells were tested for expression using both indirect immunofluorescence flow cytometry and fluorescence microscopy as previously described (Worth et al, J. Immunol. 157:5660 (1996)).

Lysosome labeling

Transfectants were grown on glass coverslips (Corning, NY) overnight at 37° C. 5 μ g of rhodamineconjugated dextran (10,000 MW, Molecular Probes, Eugene, OR) was added to each coverslip for 90 min. at 37°C . Cells were washed with PBS followed by addition of fresh media to the coverslips as described by Oh and Swanson (Oh and Swanson, J. Cell Biol. 132:585 (1996)). Imaging of lysosomes was performed using an axiovert 135 fluorescence microscope (Carl Zeiss, Thornwood, NY) utilizing mercury illumination. Optical filters for rhodamine excitation and emission were 530DF22 and 590DF30, respectively (Omega, Brattlesboro, VT). Images were 15 observed using an ICCD (Hamamatsu, Japan) coupled to a Scion LG-3 (Scion Corp., Frederick, MD) image capture board on a Dell Precision 410 Workstation (Round Rock, TX). Images were processed using Scion Image software.

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Phagocytosis of erythrocytes

SRBCs (Alsevers; Rockland Scientific, Gilbertsville, PA) were opsonized with the highest subagglutinating concentration of rabbit anti-sheep erythrocyte Ab (ICN, Costa Mesa, CA). Subsequently, antibody coated cells (EA) were added at a target-to-effector ratio of 10:1 (EA:transfectant). The EA were incubated with transfectants for 45 min. at 37° C in culture media. Coverslips were then placed on ice to stop phagocytosis.

Fluorescence microscopy

Goat anti-rabbit IgG F(ab')₂ fragments

conjugated with fluorescein isothiocyanate (ICN,

Costa Mesa, CA) were added to the coverslips for 30

min. on ice to detect the external EA. The

coverslips were observed using bright field

microscopy or by fluorescence microscopy using the

system described above. Narrow band-pass

discriminating filters were used with excitation at

482 nm and emission at 530 nm for FITC fluorescence.

Electron microscopy

Transfectants expressing either wild type FCYRIIA (131-3) or tailless FCYRIIA with CR3 (169-23) were incubated with opsonized sheep erythrocytes for 45 min. at 37°C in culture media. The cells were washed then fixed with glutaraldehyde overnight at 4°C. To detect the lysosomal compartment, acid phosphatase was stained for using modified Gormori's media consisting of 13.9mM β -glycerophosphate, 1mM $Pb(NO_3)_2$, 0.05M acetate buffer, 0.08% $CaCl_2$, and 5% sucrose. Cells were treated with the acid phosphatase stain for 1hr at 37°C with gentle agitation. The cells were washed extensively with 25 cacodylate buffer then post-fixed with osmium tetroxide for 1hr at room temperature. The cells were dehydrated and embedded in Spurr's resin as described previously (Spurr, J. Ultrastruct. Res. 26:31 (1969)). Thin-sections were viewed with a

Joel 35e (Japan) electron microscope. Micrographs were taken using an in-column digital camera system coupled to a Macintosh G3 computer and processed with Adobe photoshop 5.0.

5 Results

Receptor expression and phagocytosis Transfected CHO cells were studied for expression of FcyRIIA and CR3 utilizing flow cytometry. Several cell lines were produced. Clone 131-3 expressed wild-type FcyRIIA. 135-12 expressed the tail-minus mutant of FcyRIIA. 161-24 expressed neither of the receptors but was exposed to the transfection protocol. Clones 169-8 and 169-23 both expressed the tailless mutant FcyRIIA in 15 combination with CR3. A wild-type FcγRIIA and CR3 clone (173-46) were also constructed. Indirect immunofluorescence analysis confirmed the phenotypes of the cell lines. In addition, a phagocytosis defective FcyRIIA was utilized that had a full length cytoplasmic domain with only the tyrosine 20 residues in each of the ITAM motifs mutated to phenylalanine (FcyRIIA ITAM mutant). This mutation has previously been shown to abolish IgG-dependent phagocytosis via FcyRIIA Mitchell et al, Blood 84:1753 (1994)). FcγRIIA(ITAM mutant) was transiently transfected into untransfected CHO cells (161-30) or a CR3 expressing cell line (169-85). Expression was determined via indirect immunofluorescence quantitated by flow cytometry.

Expression of wild-type FcYRIIA and this FcyRIIA(ITAM mutant) were equivalent.

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To confirm that the receptors were functional, phagocytosis was examined using IgG-coated sheep 5 erythrocytes (EA). After incubation of EA with the transfectants for 30 min. at 37°C, it was found that the wild-type FcYRIIA (clone 131-3) was capable of internalizing IgG-coated erythrocytes. However, the FcyRIIA tailless (clone 135-12) and the FcyRIIA(ITAM mutant) (clone 161-30) were not able to phagocytose EA, as previously reported (Tuijnman et al, Blood 79:1651 (1992), Mitchell et al, Blood 84:1753 (1994), Worth et al, J. Immunol. 157:5660 (1996)). However, the co-expression of CR3 with either of the mutant FcyRIIAs (clones 169-8,169-23 and 169-85) restored FcYR-dependent phagocytosis.

Fluorescence detection of phagosome-lysosome fusion It was next determined whether the cytoplasmic tail of FcYRIIA participates in phagolysosomal 20 fusion. Fluorescently-labeled dextran was used to label lysosomes Oh and Swanson, J. Cell Biol. 132:585 (1996)). Fluorescent dextran is taken up by pinocytosis then delivered to lysosomes. This allows the fluorescent dextran to spill from the pre-loaded lysosomes into the phagosome. After incubation with dextran, the transfectants exhibited dextran located in small punctate vesicles when viewed with fluorescence microscopy.

Previous work has shown that co-expression of CR3 and a phagocytosis defective tailless FcYRIIA restored IgG-dependent phagocytosis (Worth et al, J. Immunol. 157:5660 (1996)). This approach was used, co-transfection of FcyRIIA and CR3, to examine postphagocytic events in the presence and absence of the cytoplasmic tail of Fc γ RIIA or in an ITAM mutant of FCYRIIA. Wild-type FCYRIIA (clone 131-3) transfectants exhibited co-localization of fluorescent dextran with the internalized IgG-coated 10 particle. This effect was seen as soon as 15 min. after addition of targets and did not change significantly up to 60 min. after phagocytosis. addition, more than 95% of the internalized targets were positive for lysosome fusion. However, when 15 the cell lines containing the mutant tailless form of FcyRIIA in the presence of CR3 were studied (clones 169-8 and 169-23), very little colocalization of IgG-coated cells with the dextran was observed. Little or no co-localization of 20 dextran with EA was observed from 15 min. to 60 min. after phagocytosis. Internalized targets displayed fusion with lysosomes in 6.4% and 8.7% of the cells for clones 169-8 and 169-23, respectively. These results were observed in two separate clones, suggesting consistency among similarly prepared clones. The FcYRIIA ITAM mutant (161-30) without CR3 is unable to induce phagocytosis of IgG-coated cells and therefore no lysosomal fusion can occur. However, in the presence of CR3 and FcYRIIA ITAM 30

mutant (169-85), phagocytosis was restored and near
wild type levels of lysosome fusion was detected.
Clone 173-46, which expressed wild-type FcγRIIA and
CR3, to determine if CR3 might affect phagolysosome
formation. Expression of CR3 did not affect the
ability of wild-type FcγRIIA to participate in
phagolysosome fusion.

Electron microscopy of phagosome-lysosome fusion As a second independent means of detecting 10 phagosome-lysosome fusion following phagocytosis, electron microscopy was employed using a specific lysosomal stain. Acid phosphatase is an enzyme specific for lysosomes and has been used extensively to stain CHO cells (Gennaro et al, Proc. Soc. Exp. 15 Biol. Med. 198:591 (1991)). Therefore, this enzyme was used to detect the localization of lysosomal enzymes inside cells. After incubation of transfectants expressing either wild type FcYRIIA or tailless FcyRIIA in the presence of CR3 with IgGcoated sheep erythrocytes, the cells were fixed and stained for acid phosphatase. After embedding, thin sections were viewed with an electron microscope. Acid phosphatase appeared as dark electron dense 25 patches, revealing the location of lysosomal enzyme activity. In the presence of the wild-type FcyRIIA (clone 131-3) acid phosphatase staining was observed near the internalized target, indicating phagolysosomal fusion. However, cells expressing 30 the tail-minus form of FcyRIIA (clone 169-8) did not

support phagolysosome formation. Thus, the acid phosphatase staining was found throughout the entire cytoplasm as punctate granules and was not localized near internalized targets. These results suggest that the cytoplasmic domain of FcyRIIA targets the internalized particle for fusion with lysosomes. The data demonstrate that the cytoplasmic tail of FCYRIIA participates in phagolysosomal fusion and that this signal is distinct from a functional ITAM.

EXAMPLE 2 10

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Lysosomal Fusion Following FcYRIIA Phagocytosis is Mediated by an L-T-L Motif

This study was designed to elucidate the mechanism by which FcyRIIA mediates lysosomal fusion. As indicated in Example 1, a mutant FcyRIIA lacking a cytoplasmic domain is not able to mediate phagocytosis. However, the presence of complement receptor type 3 (CR3) restores phagocytosis, but no lysosomal fusion is observed. Therefore, the 20 cytoplasmic domain of FcγRIIA is required for lysosomal fusion. The FcYRIIA cytoplasmic domain ITAM (immunoreceptor tyrosine-based activation motif) was disabled to determine if an intact ITAM is required for lysosomal targeting. Mutation of both tyrosines in the ITAM to phenylalanine abolished phagocytosis. However, co-transfection of CR3 with this ITAM mutant restored phagocytosis and wild-type (WT) levels of lysosomal fusion were

observed. After mutation of signaling sequences in the cytoplasmic domain of FcyRIIA, it was noted that a novel L-T-L motif at the C-terminal of the ITAM was responsible for targeting of FcyRIIA 5 internalized targets to the lysosomal compartment, but not required for the initial stage(s) of phagocytosis. Mutation of either of the leucine residues individually or in tandem resulted in 70% (p<0.05 compared to wt FcyRIIA) inhibition of internalized targets to co-localize with lysosomes pre-loaded with fluorescent dextran. Mutation of the threonine alone elicited similar results, thus abolishing 78% (p<0.05 compared to wt FcyRIIA) of co-localization. However, when the L-T-L motif was 15 mutated to A-A-A, lysosomal targeting was abolished as observed with tailless FcyRIIA. Therefore, a novel L-T-L motif in the cytoplasmic domain of FCYRIIA is responsible for mediating phagolysosomal fusion. (See also Fig. 1).

20 EXAMPLE 3

FCYRIIA wild-type (IIA), various mutants of the L_T-L motif in the cytoplasmic domain of FCYRIIA (IIA(YLTA), IIA(YATL), IIA(YATA), IIA(YAAA)), or FCYRIIA lacking a cytoplasmic domain (IIA(tailless)) were transfected into chinese hamster ovary (CHO) cells. These cells were pre-loaded with fluorescently labeled dextran by incubating the cells with medium containing TRITC-dextran. The

cells were then allowed to phagocytose IgG-coated erythrocytes (EA) for 30 minutes. After 30 minutes the cells were placed on ice to stop phagocytosis and observed for location of the internalized EA and TRITC-dextran. Data presented in Fig. 2 are shown as percent of internalized EA colocalized with TRITC-dextran. As shown, mutation of the L-T-L motif inhibits the colocalization (phagolysosome fusion) of the internal EA with TRITC-dextran.

The data presented in Fig. 3 demonstrate that the L-T-L motif mediates specific targeting of internalized targets to fuse with lysosomes. In time-course experiments, the mutant FcYRIIA containing a mutant L-T-L motif, inhibited phagolysosome formation at early time points compared to wild-type FcYRIIA.

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To elucidate the mechanism by whih the L-T-L motif inhibits phagolysosome fusion, another marker of lysosome location was studied. Lysosome associated membrane protein (LAMP) is a cytosolic protein that colocalizes with lysosomes and the plasma membrane. It was observed that the L-T-L mutation inhibits the spilling of fluorescent dextran into phagosomes but does not inhibit the acquisition of lysosome associated proteins thus suggesting that phagolysosome formation may be a more complex process than originally thought (see Fig. 4).

The common γ-chain does not mediate efficient phagolysosome fusion. A chimeric molecule was

produced containing the ligand-binding domain of
FcγRIII and the γ-chain transmembrane and
cytoplasmic domain. Upon insertion of the L-T-L
motif into the cytoplasmic domain of the chimeric
molecule, a 50% increase in phagolysosome formation
was observed (see Fig. 5). These data indicate that
insertion of the L-T-L motif into a receptor that is
not efficient in mediating phagolysosomal fusion can
be used to increase the ability of receptors to kill
bacterium

All documents cited above are hereby incorporated in their entirety by reference.

WHAT IS CLAIMED IS:

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A method of mediating trafficking of a target to a lysosomal compartment of a cell comprising associating with said target a molecule containing an L-T-L peptide under conditions such that said trafficking is effected, wherein said molecule is other than an FcγRIIA molecule endogenous to said cell.

- The method according to claim 1 wherein
 said target is a bacterium.
- A method of enhancing the ability of a cell to degrade a particle comprising introducing into said cell a nucleic acid sequence encoding an Fc receptor comprising an L-T-L sequence under conditions such that said nucleic acid sequence is expressed and said enhancement is effected.
 - 4. The method according to claim 3 wherein said Fc receptor is FcyRIIA or a modified form thereof comprising at least 1 additional L-T-L peptide in the cytoplasmic domain thereof.
 - 5. The method according to claim 3 wherein said Fc receptor comprises a γ chain comprising at least one L-T-L peptide in the cytoplasmic domain thereof.

 The method according to claim 3 wherein said cell naturally expresses FcγRIIA.

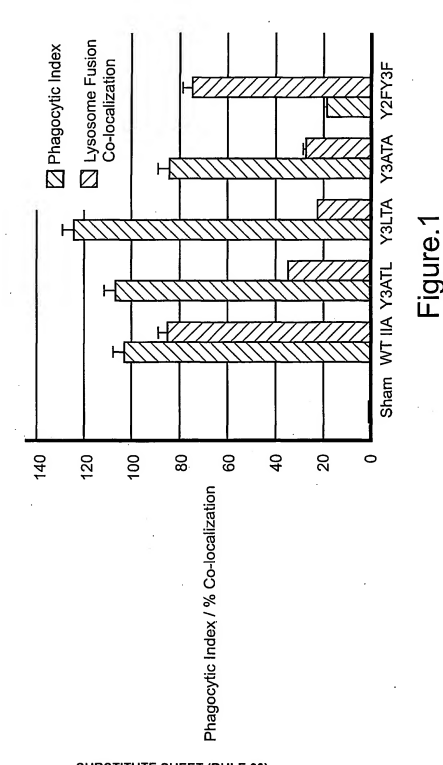
- 7. The method according to claim 3 wherein said cell does not naturally express FcyRTIA.
- 5 .8. The method according to claim 3 wherein said cell is an endothelial cell, a fibroblast, a macrophage or an epithelial cell.
 - 9. The method according to claim 3 wherein said particle is a bacterium.
- 10. The method according to claim 3 wherein said nucleic acid sequence is introduced into said cell in a liposome, a bacterium or a viral vector.
 - 11. A method of enhancing the ability of a cell that has on its surface FcyRIIA to degrade a particle comprising contacting said cell with an agent that increases the number of FcyRIIA molecules present on the surface of said cell so that said enhancement is effected.
- 12. The method according to claim 11 wherein said cell is a macrophage.
 - 13. The method according to claim 11 wherein said agent is IFN- γ or an inhibitor of IL-4.

14. A method of targeting a particle to an FcyRIIA molecule present on a cell comprising contacting said particle with a bi-specific antibody that recognizes said particle and the extracellular domain of said FcyRIIA under conditions such that said antibody binds said particle and said FcyRIIA so that said targeting is effected.

- 15. The method according to claim 14 wherein said particle is a microorganism.
- 16. A method of inducing uptake and targeting of a particle to a phagolysosome of a cell comprising contacting said particle with an IgG antibody that recognizes said particle, which antibody has associated therewith an L-T-L- containing peptide, wherein said contacting is effected under conditions such that said antibody binds said particle and said induction is effected.
 - 17. The method according to claim 16 wherein said particle is an immune complex.
- 20 18. The method according to claim 16 wherein said particle is a microorganism.
 - 19. A method of inducing targeting of a particle to a phagolysosome of a cell comprising contacting said particle with an L-T-L-containing

peptide, wherein said contacting is effected under conditions such that said an L-T-L-containing peptide associates with said particle and said induction is thereby effected.

- 20. A method of inhibiting degradation of a particle by a phagolysosome of a cell comprising contacting said cell with an agent that inhibits the function of L-T-L-containing molecule present in said cell so that said inhibition is effected.
- 10 21. The method according to claim 2 wherein said bacterium is a mycobacterium or an antibiotic resistant bacterium.
- 22. The method according to claim 9 wherein said bacterium is a mycobacterium or an antibiotic resistant bacterium.
 - 23. The method according to claim 15 wherein said microorganism is a mycobacterium or an antibiotic resistant bacterium.
- 24. The method according to claim 18 wherein said microorganism is a mycobacterium or an antibiotic resistant bacterium.
 - 25. The method according to claim 20 wherein said particle is an IgG-coated immune complex.



SUBSTITUTE SHEET (RULE 26)

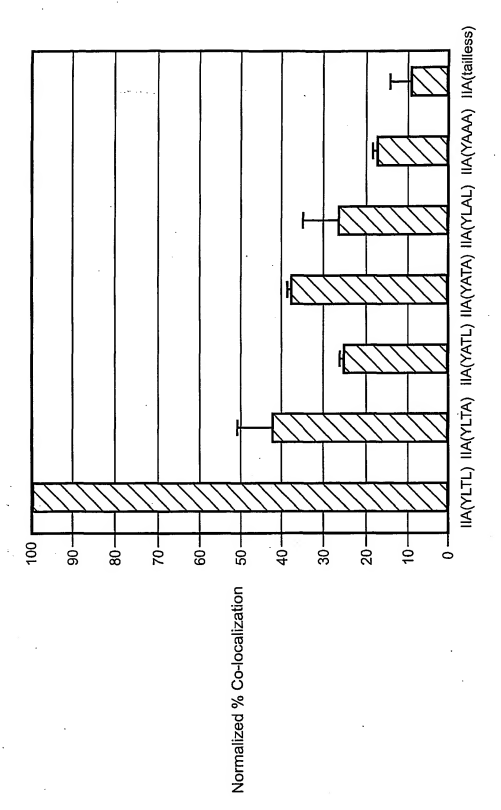
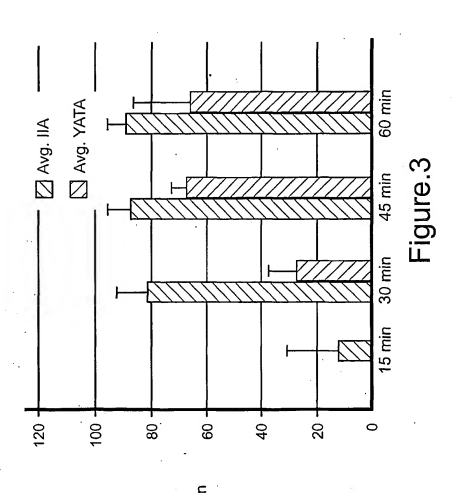
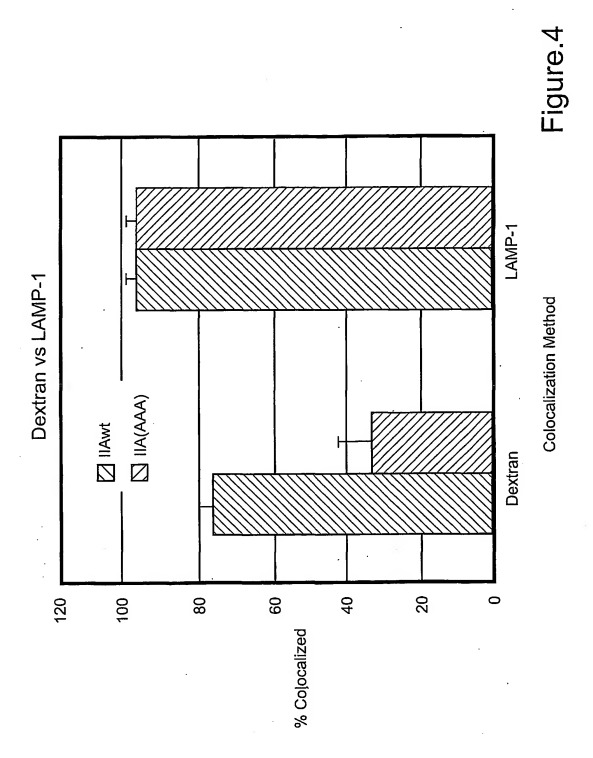
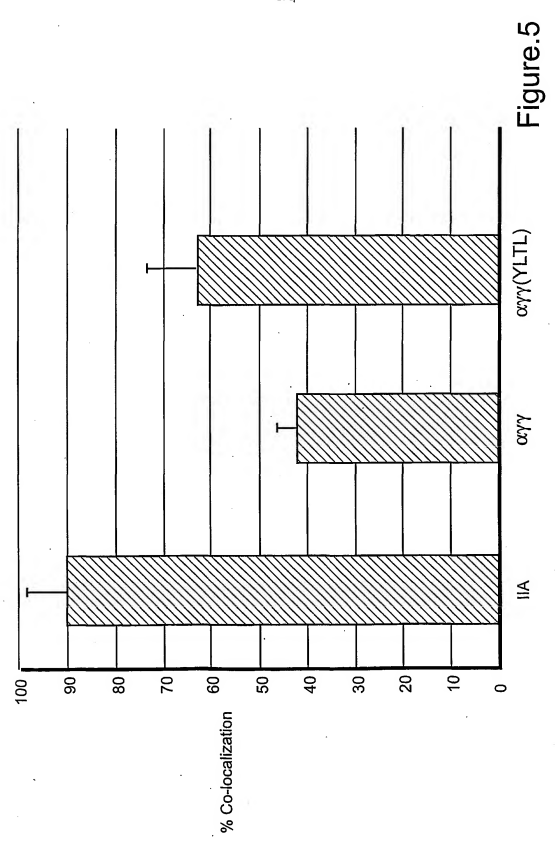


Figure.2



% Colocalization





SUBSTITUTE SHEET (RULE 26)

International application No.

PCT/US01/43340

		FC1763017433	-10	
A. CLASSIFICATION OF SUBJECT MATTER				
IPC(7) : G01N 33/53				
US CL : 435/7.1				
According to International Patent Classification (IPC) or to both national classification and IPC				
B. FIELDS SEARCHED				
Minimum documentation searched (classification system followed by classification symbols)				
U.S.: 435/7.1				
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Documentati	on searched other than minimum documentation to the	e extent that such documents are inclu-	led in the fields searched	
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	ata base consulted during the international search (nar	ne of data base and, where practicable	, search terms used)	
Please See Continuation Sheet				
C. DOCUMENTS CONSIDERED TO BE RELEVANT				
Category *	Citation of document, with indication, where ap	propriete of the relevant passages	Relevant to claim No.	
				
Х	DOWNEY, G.P. et al. Phagosomal Maturation, Ac Growth in Nonphagocytic Cells Transfected with Po		1-2	
	Biological Chemistry. 01 October 1999, Vol 274,	No 40 pages 28426 28444	1	
	especially pages 28436 and 28443.	110. 40, pages 20430-20444,	1	
	especially pages 20450 and 20445.		l f	
Y	ODTAIO MACNE at al. Identification of the Surf	one Expected Lipids on the Cell	21	
1	ORTALO-MAGNE et al. Identification of the Surf Envelopes of Mycobacterium tuberculosis and Othe		21 .	
	Bacteriology. January 1996, Vol. 178, No. 2, page		1	
	Bacteriology. January 1990, Vol. 178, 180. 2, page	as 450-401, especially page 450.	l	
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	demonstration of Dec C	San anti-of-		
	r documents are listed in the continuation of Box C.	See patent family annex.		
Special categories of cited documents: "T" later document published after the international filing date or prio date and not in conflict with the application but cited to understan				
"A" document	t defining the general state of the art which is not considered to be	principle or theory underlying the i		
of particular relevance				
"B" carlier ap	oplication or patent published on or after the international filing date		idered to involve an inventive step	
	when the document is taken alone			
"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as "Y" document of particular relevance; the claimed invention cannot be			he claimed invention cannot be	
specified)		considered to involve an inventive	step when the document is	
"O" document referring to an oral disclosure, use, exhibition or other means		combined with one or more other a being obvious to a person skilled in	uch documents, such combination	
"O" document referring to an oral disclosure, use, exhibition or other means		•		
"P" document published prior to the international filing date but later than the "&" document member of the same patent family				
priority date claimed				
Date of the	Date of the actual completion of the international search Date of mailing of the international search report			
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04 March 2002 (04.03.2002) Property of the state of the				
Name and mailing address of the ISA/US Authorized officer Authorized officer				
Commissioner of Patents and Trademarks Box PCT Margaret Jannoz Jaw Servel Jordan			tor	
Washington, D.C. 20231				
Facsimile No	Facsimile No. (703) 305-3230 Telephone No. (703) 308-0196			
D. DOWNER MAD C. L. L. C. C. L. COO.				

Form PCT/ISA/210 (second sheet) (July 1998)

International application No.

PCT/US01/43340

constitute a special technical feature within the meaning of PCT Rule 13.2 and that each of such methods accordingly defines a separate invention.			
separate invention.			
Continuation of B. FIELDS SEARCHED Item 3:			
WEST, PubMed traffick, phagocytosis, lysosome, FcgRIIA, bacteria, mycobacteria, capsule, inventor name search			
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Form PCT/ISA/210 (second sheet) (July 1998)

International application No.

PCT/US01/43340

Box I Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)			
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:			
Claim Nos.: because they relate to subject matter not required to be searched by this Authority, namely:			
2. Claim Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:			
3. Claim Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).			
Box II Observations where unity of invention is lacking (Continuation of Item 2 of first sheet)			
This International Searching Authority found multiple inventions in this international application, as follows: Please See Continuation Sheet			
As all required additional search fees were timely paid by the applicant, this international search report covers all			
searchable claims.			
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.			
3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:			
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-2 and 21			
Remark on Protest The additional search fees were accompanied by the applicant's protest.			
No protest accompanied the payment of additional search fees.			

International application No.

PCT/US01/43340

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING

This application contains the following inventions or groups of inventions which are not so linked as to form a single general inventive concept under PCT Rule 13.1. In order for all inventions to be examined, the appropriate additional examination fees must be paid.

Group 1, claim(s) 1-2 and 21, drawn to a method of mediating trafficking of a target to a lysosomal compartment of a cell comprising associating with said target a molecule containing an LTL peptide.

Group 2, claim(s) 3-10 and 22, drawn to a method of enhancing the ability of a cell to degrade a particle comprising administering a nucleic acid sequence encoding an Fc receptor comprising an LTL sequence.

Group 3, claim(s) 11-13, drawn to a method of enhancing the ability of a cell to degrade a particle comprising contacting the cell with an agent that increases the number of FogammaRIIA molecules, wherein the agent is IFN-gamma.

Group 4, claim(s) 11-13, drawn to a method of enhancing the ability of a cell to degrade a particle comprising contacting the cell with an agent that increases the number of FogammaRIIA molecules, wherein the agent is an inhibitor of IL-4.

Group 5, claim(s) 14-15 and 23, drawn to a method of targeting a particle to a FegammaRIIA molecule comprising contacting a particle with a bi-specific antibody, and administering the particle/antibody complex.

Group 6, claim(s) 16-17, drawn to a method of inducing uptake and targeting of a particle to a phagolysosome of a cell comprising contacting said particle with an IgG antibody associated with an LTL containing peptide, wherein the particle is an immune complex.

Group 7, claim(s) 16, 18, and 24, drawn to a method of inducing uptake and targeting of a particle to a phagolysosome of a cell comprising contacting said particle with an IgG antibody associated with an LTL containing peptide, wherein the particle is a microorganism.

Group 8, claim(s) 19, drawn to a method of inducing targeting of a particle to a phagolysosome of a cell comprising contacting said particle with an LTL containing peptide.

Group 9, claim(s) 20 and 25, drawn to a method of inhibiting degradation of a particle by a phagolysosome of a cell comprising contacting said cell with an agent that inhibits the function of LTL containing molecules present in the cell.

The inventions listed as Groups 1-9 do not relate to a single general inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons:

Claims 1-2 lack a special technical feature over Downey et al. (J. Biol. Chem. 1999; 274(40): 28436-28444). Downey et al. teach a method of mediating trafficking of a target (E. coli) to a lysosomal compartment of a cell comprising associating with said target (E. coli) a molecule containing an LTL peptide under conditions such that trafficking is effected, wherein said molecule is other than an FcgRIIA molecule endogenous to said cell (i.e. a CHO cell transfected with FcgRIIA; see pages 28436 and 28443 in particular). The instant specification discloses that FcgRIIA has an LTL peptide that mediates phagocytosis.

Claim 21 lacks a special technical feature over Downey et al. (J. Biol. Chem. 1999; 274(40): 28436-28444) and Ortalo-Magne et al. (J. of Bacteriology 1996; 178(2): 456-461). Downey et al. has been discussed supra. Ortalo-Magne et al. teach that Mycobacterium have a capsule (see page 456 in particular). It would have been obvious to substitute the Mycobacterium taught by Ortalo-Magne et al. for the E. coli taught by Downey et al. because both types of bacteria have capsules that would be recognized by FcgRIIA.

Pursuant to 37 C.F.R. 1.475(d), the ISA/US considers that where multiple products and processes are claimed, the main invention shall consist of the first invention of the category first mentioned in the claims, and the first recited invention of each of the other categories related thereto. Accordingly, the main invention (Group I) comprises the first method. Further pursuant to 37 C.F.R. 1.475(d), the ISA/US considers that any feature which the subsequently recited methods share with the main invention does not